Ref #	Hits	Search Query	DBs	Default Operator	Plurals	Time Stamp
L1	1	("6034029").PN.	US-PGPUB; USPAT; USOCR	OR	OFF	2005/04/28 17:26
L2	1	("5354915").PN.	US-PGPUB; USPAT; USOCR	OR	OFF	2005/04/28 17:27
L3	4859	Zirconium adj dioxide	US-PGPUB; USPAT; USOCR	OR	ON	2005/04/28 17:27
L4,	4198	monoclinic	US-PGPUB; USPAT; USOCR	OR	ON	2005/04/28 17:27
L5	7045	tetragonal	US-PGPUB; USPAT; USOCR	OR	ON	2005/04/28 17:27
L6	155738	cubic	US-PGPUB; USPAT; USOCR	OR	ON	2005/04/28 17:28
L7	161998	14 or 15 or 16	US-PGPUB; USPAT; USOCR	OR	ON	2005/04/28 17:28
L8	710	13 and 17	US-PGPUB; USPAT; USOCR	OR	ON	2005/04/28 17:28

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                 data from INPADOC
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     13 MAR 22 REGISTRY/ZREGISTRY enhanced with experimental property tags
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                 fields
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     15 APR 04
                EMBASE - Database reloaded and enhanced
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     17 APR 25
                Patent searching, including current-awareness alerts (SDIs),
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NEWS
      18 APR 28
                Improved searching of U.S. Patent Classifications for
                 U.S. patent records in CA/CAplus
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This file contains CAS Registry Numbers for easy and accurate substance identification.

=> s "zirconium dioxide"

188973 "ZIRCONIUM"

21 "ZIRCONIUMS"

188976 "ZIRCONIUM"

("ZIRCONIUM" OR "ZIRCONIUMS")

430276 "DIOXIDE"

6427 "DIOXIDES"

431897 "DIOXIDE"

("DIOXIDE" OR "DIOXIDES")

8067 "ZIRCONIUM DIOXIDE"

("ZIRCONIUM"(W) "DIOXIDE")

=> s monoclinic

L1

83888 MONOCLINIC

4 MONOCLINICS

83891 MONOCLINIC

(MONOCLINIC OR MONOCLINICS)

128 MONOCLIN

L2 84011 MONOCLINIC

(MONOCLINIC OR MONOCLIN)

=> s tetragonal

49831 TETRAGONAL

7 TETRAGONALS

L3 49834 TETRAGONAL

(TETRAGONAL OR TETRAGONALS)

=> s cubic

91524 CUBIC 22 CUBICS

L4 91531 CUBIC

(CUBIC OR CUBICS)

=> s 12 or 13 or 14

L5 207262 L2 OR L3 OR L4

=> s 11 and 15

L6 1135 L1 AND L5

=> s catalyst

686360 CATALYST

690197 CATALYSTS

L7 879924 CATALYST

(CATALYST OR CATALYSTS)

=> s 16 and 17

L8 41 L6 AND L7

=> s hydrogen?

L9. 1116688 HYDROGEN?

 \Rightarrow s 18 and 19

L10 8 L8 AND L9

=> d 110 1-8 abs ibib

ANSWER 1 OF 8 CAPLUS COPYRIGHT 2005 ACS on STN
Gallium-promoted sulfated zirconia (GSZ) catalysts were prepared
by impregnation of zirconium hydroxide with aqueous Ga2(SO4)3 followed by
calcination. Isomerization of n-hexane was studied over GSZ at
150', 2.0 MP, WHSV 2 and H2/hexane (molar) ratio of 5. In
comparison to sulfated zirconia (GSZ) was greatly improved and it
remained stable at 85t. In particular, almost all the products were
isomers of hexane and the selectivity of 2,2-MB reached 201. The results
of characterization indicated that the addition of gallium onto SZ
catalyst showed little difference in acid strength between SZ and
GSZ catalysts while the redox properties of the SZ
catalyst changed with addition of gallium. The transformation of SZ
crystalline from metastable tetragonal phase, the more active phase,
to smoodinic phase was retarded with the addition of gallium.
Also, the simultaneous promotion of Pt and Ga brings the production
distribution very close to the equilibrium one.
ESSION NUMBER:
UNENT NUMBER:
140:G1930
Hydroisomerization of n-hexane over gallium-promoted
sulfated zirconia
Cao, Chong-jiang Han, Song Chen, Chang-Lin; Xu,
Nan-Ping, Mou, Chuny-Yuan
College of Chemistry and Chemical Engineering, Nanjing
University of Technology, Nanjing, 210009, Peop. Rep.
China
CRCE:

Catalysis Communications (2003), 4(10), 511-515

ACCESSION NUMBER: DOCUMENT NUMBER: TITLE:

AUTHOR (S):

CORPORATE SOURCE:

China
Catalysis Communications (2003), 4(10), 511-515
CODEN: CCAOAC: ISSN: 1566-7367
Elsevier Science B.V.

SOURCE:

PUBLISHER: DOCUMENT TYPE: LANGUAGE: REFERENCE COUNT: Journal English

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ANSWER 3 OF 8 CAPLUS COPYRIGHT 2005 ACS on STN
The activity and durability of the catalysts prepared by the oxidation-reduction treatment of amorphous Co-15 atomics Zr, Ni-40 atomics

AB The activity and durability of the catalysts prepared by the oxidation-reduction treatment of amorphous Co-15 atomic 2r, Ni-40 atomic 2r, and
Ni-30 atomic 2r-10 atomic 3r alloys were investigated for simultaneous methanation of carbon monoxide and carbon dioxide. The Ni-302r-105m catalyst showed the highest activity among the catalysts examined; the activity of the Co-152r catalyst is lower than those of the nickel-based catalysts, in agreement with the activity of the solo methanation of carbon dioxide. On all the catalysts, carbon monoxide reacts preferentially with hydrogen and is completely converted into methane at 2523 K. The remaining hydrogen further reacts with carbon dioxide to form methane. The methanation rate in the H2-CO-CO2 mixed gas was higher than that in H2-CO mixed gas without CO2. This is probably related to the prevention of the formation of surface carbon by disproportionation of carbon monoxide due to the presence of carbon dioxide. The activity of the Ni-402r catalyst at 573 K yradually decreased with reaction time.

Tetragonal 2ro2, the presence of which is responsible for the high activity, is transformed to thermodynamically more stable monoclinic 2ro2 during the reaction. In contrast to the Ni-402r catalyst, the Ni-302r-105m catalyst sustains the initial high activity, and no structural changes were observed during the durability test regardless of the presence of a small amount of H2S.

ACCESSION NUMBER: 1998:571532 CAPLUS

DOCUMENT NUMBER: 1998:571532 CAPLUS

CORPORATE SOURCE: Institute for Materials Research, Tohoku University, Sendai, 980-8577, Japan

SOURCE: Applied Catalysis, A: General (1998), 172(1), 131-140 CODER: ACACE4; 15SN: 0926-860X

FUBLISHER: Slevier Science B.V.

JOURNAL THERE ARE 13 CITED REFERENCES AVAILABLE FOR THIS

Journal English 13 Ti

PUBLISHER: DOCUMENT TYPE: LANGUAGE: REFERENCE COUNT:

THERE ARE 13 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

ANSWER 2 OF 8 CAPLUS COPYRIGHT 2005 ACS on STN
Zirconia-loaded alumina samples were prepared and used as supports for
platinum and tin metallic phases. X-ray diffraction patterns of 9,0 and
13.0 wt \$2\text{r02-loaded alumina samples showed the characteristic lines of
the tetragonal ZrO2 phase. For the base alumina and these two
supports, the nitrogen adsorption-desorption isotherms displayed type IV
isotherms and a type HI hysteresis loop characteristic of mesoporous
materials, with EET areas and pore vols. decreasing with increasing ZrO2
content. Platinum and tin were incorporated into these substrates and
then characterized by temperature-programmed reduction, XPS, and their
formance

ormance in regard to n-butane dehydrogenation. The TPR profiles of the bimetallic systems revealed that platinum is readily reduced whereas tin reduction depends on the support, the metal loading, the preparation method, and the pretreatment conditions. Upon hydrogenation up to 773 K, tin undergoes partial reduction to the metal and the SnO/Sn2+ ratio increases

undergoes partial reduction to the metal and the SnO/Sn2+ ratio increases with increasing ZrO2 content. All these systems were highly selective toward olefins (i.e., n-butenes), with a minor contribution of the isomerization and cracking reactions. The catalysts became deactivated by coke deposition, although this deactivation was less marked in the Pt-Sn deposited on the ZrO2-Al2O3 substrates.

ACCESSION NUMBER: 2000:83508 CAPLUS DOCUMENT NUMBER: 134:133906
ITILE: Alumina- and Zirconia-Alumina-Loaded Tin-Platinum. Surface Features and Performance for Butane Dehydrogenation

AUTHOR(S): Largese C.I Campos-Martin, J. M.) Fierro, J. L. G. CORPORATE SOURCE: Inst. Catalisis Petroleoquimica, CSIC, Madrid, 28049, Spain

SOURCE: Langmuir (2000), 16(26), 10294-10300 CODE: Langmuir (2000), 16(26), 10294-7463

PUBLISHER: Aperican Chemical Society Journal English

REFERENCE COUNT: REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE REFORMAT

THERE ARE 46 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L10 ANSWER 4 OF 8 CAPLUS COPYRIGHT 2005 ACS on STN

AB 2r02, which is 280 weight monoclinic and has a surface area of 2100 m2/q, is produced by reaction of aqueous Zr salt solns. with NHB, aging to convert a tetragonal phase to a monoclinic phase at 0-300°, drying, and calcining at 200-600°. The product is suitable as a catalyst or catalyst support, especially for hydrogenation, dehydrogenation, Fischer-Tropsch syntheses, desulfurization, isomerization, polymerization, and steam reforming.

ACCESSION NUMBER: 1998:410662 CAPLUS

TITLE: Monoclinic sirconium dioxide with high surface area ...c xirconium
.xide with high sur
Wulff-Doring, Joachim;
Ferdi
BASF A.-G., Germany
Eur. Pat. Appl., 9 pp.
CODEN: EFXXDW
Patent
German
1 dioxide with high surface area Wulff-Doring, Joachim; Stichert, Wolfram; Schuth, INVENTOR(S): PATENT ASSIGNEE(S): SOURCE: DOCUMENT TYPE: LANGUAGE: FAMILY ACC. NUM. COUNT: PATENT INFORMATION: PATENT NO. KIND DATE APPLICATION NO.

PAILENT NO.

EP 849224
B1 20010613
R: AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, LU, NL, SE, MC, PT, IE, SI, LT, LV, FI, RO

DE 19653629
A1 19980625
DE 1996-19653629
JA 19980625
DE 1996-19653629
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DE 1996-19653629
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DE 1996-19653629
JA 19981217
DE 1996-19653629
JA 19981218
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DE 1996-19653629
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DE 1996-19653629
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DE 1996-19653629
A 19961200
DE 19961200 I, RO
19980625 DE 1996-19653629 19961220
20000307 US 1997-992569 19971217
20011016 ES 1997-122267 19971217
19981007 CN 1997-107288 19971219
19981215 JF 1997-352888 19971222
DE 1996-19653629 A 19961220
THERE ARE 6 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT ANSWER 5 OF 8 CAPLUS COPYRIGHT 2005 ACS on STN Pure monoclinic and tetragonal sirconium dioxides were successfully prepared at pH values of .apprx.9.5 and 211.5, rep. Their crystal structures were characterized by x-ray diffraction. Calcination temperature has a great influence upon crystals.

when calcination temperature was a great instruence upon crystals.
calcination temperature varied from 350' to 500', amorphous ZrO2 converted to crystal phase and the amount of monocilinic phase increased, while that of tetragonal phase decreased. Different crystal phases have different catalytic performance. Monocilinic ZrO2 catalyst has a higher selectivity for isobutene in F-T synthesis, while the amount of isobutene vas reduced over tetragonal ZrO2. Catalytic properties of ZrO2 prepared by using supercrit, fluid drying method is better than those by other methods. Reaction mechanisms are discussed.

ACCESSION NUMBER: 1997:624568 CAPLUS 127:263082

TITLE: Isobutene formation from synthesis gas over irronium dioxide

AUTHOR(S): CORPORATE SOURCE:

1997:624568 CAPLUS
127:263082
1sobutene formation from synthesis gas over
zirconium dioxide
Wang, Guojun: Su, Guiqin: Yin, Yuanqi
Chinese Academy Sciences, Lanzhou Inst. Chemical
Physics, Lanzhou, 730000, Peop. Rep. China
Fenzi Cuihua (1997), 11(4), 278-282
CODEN: FECUEN: ISSN: 1001-3555
Zhongquo Kexueyuan Lanzhou Huaxue Wuli Yanjiuso
Journal

SOURCE:

PUBLISHER: DOCUMENT TYPE: LANGUAGE:

DOCUMENT TYPE: LANGUAGE: FAMILY ACC. NUM. COUNT: PATENT INFORMATION:

KIND DATE APPLICATION OF THE PROPERTY OF THE P PATENT NO. EP 607591 EP 607591 EP 607591 EP 607591 R: AT, BE, CH, US 5354915 JP 06226093 JP 2864089 AT 185091 ES 2136110 PRIORITY APPLN. INFO.: OTHER SOURCE(S):

ANSWER 8 OF 8 CAPLUS COPYRIGHT 2005 ACS on STN
Metal-support interactions are proposed to explain a much lower
chemisorption of H on Rh/ZrO2 compared to Rh/Y-Al203
catalysts. The decrease in H Chemisorption, while retaining the
ability to chemisorb O, is a characteristic of strong metal-support
interactions. Well dispersed samples of Rh203 on ZrO2 were prepared and the
structural properties of the catalysts were related to their
stability in reducing atmospheres. Changes in the structure of the phases
formed and the temps. at which reduction to Rh metal occurred were then
related to the relative strengths of the catalyst-support
interactions. A comparison of the stability towards reduction of the bulk dispersed Rh203 demonstrates the influence of an interaction between the dispersed metal oxide and the support.

ACCESSION NUMBER: 1987:39138 CAPLUS
DOCUMENT NUMBER: 106:39138
Preparation and characterization of dispersed rhodium 106:39138
Preparation and characterization of dispersed rhodium oxide on tetragonal zirconium oxide
Zhang, Y. C.; Dwight, K.; Wold, Aaron
Dep. Chem., Brown Univ., Providence, RI, USA
Report (1986), TR-39; Order No. AD-A166345/9/GAR, 8
pp. Avail: NTIS
From: Gov. Rep. Ahnounce. Index (U. S.) 1986, 86(15),
Abstr. No. 633,336
Report
English AUTHOR(S): CORPORATE SOURCE: SOURCE: DOCUMENT TYPE: LANGUAGE:

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